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Sustainable wildlife extraction and the impacts of socio-economic change among the
Kukama-Kukamilla people of the Pacaya-Samiria National Reserve, Peru.

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Abstract

Throughout the tropics, hunting and fishing are critical livelihood activities for many Indigenous peoples. However, these practices may not be sustainable following recent socio-economic changes in Indigenous populations. Aiming to understand how human population growth and increased market integration affect hunting and fishing patterns, we conducted semi-structured interviews in five Kukama-Kukamilla communities living along the boundary of the Pacaya-Samiria National Reserve, in the Peruvian Amazon. Extrapolated annual harvest rates of fish and game species by these communities amounted to 1,740 t and 4,275 individuals (67 t), respectively. At least 23 fish and 27 game species were harvested. We found a positive correlation between village size and annual total community-level harvest rates of fish and a negative relationship between market exposure and mean per-capita harvest rates of fish. Catch-per-unit-effort (CPUE) analyses indicated local depletion of fish populations around larger, more commercial communities. CPUE of fish was lower in more commercial communities and fishermen from the largest village travelled farther into the reserve, where CPUE was higher. We found no effect of village size or market exposure on harvest rates or CPUE of game species. However, larger, more commercial communities targeted larger, economically valuable species. This study provides evidence that human population growth and market-driven hunting and fishing pose a growing threat to wildlife and Indigenous livelihoods through increased harvest rates and selective harvesting of vulnerable species.

Keywords: Sustainability; Hunting; Fishing; Protected area; Amazon.

Introduction

In tropical forests, hunting and fishing are crucial to the livelihoods of Indigenous peoples as a source of protein and income (East et al., 2005). Unfortunately, a growing number of studies suggest current harvests of a variety of species exceed sustainable levels, causing widespread population declines and local extinctions (Abernethy et al., 2013; Castello et al., 2014; Morcatty & Valsecchi, 2015; Parry & Peres, 2015). As a result, the sustainability of hunting and fishing has become the subject of considerable concern among ecologists, anthropologists, protected area managers and conservationists alike. This has sparked a debate surrounding the presence of Indigenous peoples in protected areas, between those who view them as a direct threat to biodiversity and those who view them as conservation allies (da Silva et al., 2005; Ohl-Schacherer et al., 2007). In-depth monitoring of hunting and fishing is a key prerequisite to promoting the sustainable use of natural resources, avoiding extinctions of important species while preserving the rights of Indigenous peoples to land, traditions, and culture.

The decreasing sustainability of hunting and fishing practices has been attributed in part to the rapid growth in Indigenous populations and their integration into the market economy. These trends have triggered powerful socio-economic changes, leading to an increasing demand for wildlife products from both the rural and urban populations and a growing economic incentive to hunt and fish commercially

(McSweeney & Jockisch, 2007; Ohl-Schacherer et al., 2007; Suarez et al., 2009; Fa et al., 2015). Simultaneously, improved technologies and transportation have enhanced the capacity of a growing number of hunters and fishermen to capture prey, including in previously inaccessible areas (Wilkie et al., 2000; Godoy et al., 2010; Foerster et al., 2012). Yet, empirical studies have revealed mixed and even positive effects of socio-economic development on wildlife harvesting (Lu, 2007). For example, opportunities for permanent and well-paid jobs combined with a preference among wealthier households for alternative protein sources like store-purchased meat can lead to a reduction in wildlife harvesting (Wilkie & Godoy, 2001; Gray et al., 2015; Vasco & Sirén, 2016). Understanding the complex interactions between socio-economic factors and extractive activities in a variety of social, cultural, and natural contexts remains imperative, especially given the need to alleviate poverty among Indigenous peoples.

In the Peruvian Amazon, hunting and fishing constitute integral components of the Kukama-Kukamilla culture. This Indigenous group harvests a large variety of natural resources from their surrounding areas that include the Pacaya-Samiria National Reserve (PSNR). In the past, a strict protectionist system in this reserve provoked a backlash of rampant poaching and over-exploitation by the local people (Bodmer et al., 2008). In the late 1990s, a new reserve administration adopted a co-management approach, permitting low levels of hunting and fishing by the local people. Since then, populations of key species have been increasing in the reserve, including threatened species such as the woolly monkey *Lagothrix* spp., lowland tapir *Tapirus terrestris* and paiche *Arapaima gigas* (Bodmer & Puertas, 2007). However, like many other Amazonian communities, the Kukama-Kukamilla are undergoing rapid socio-economic changes that could once again increase pressure on wildlife.

In this study, we aimed to explore how socio-economic factors influence the hunting and fishing patterns of the Kukama-Kukamilla people. The results of this study provide important insights into the factors that underpin sustainable resource use, specifically the risk of human population growth and market-driven hunting and fishing brought about by rural development. Previous studies have generally explored the effects of socio-economic conditions on wildlife harvesting between households. However, because households within a community harvest wildlife from a communal catchment area, we explored the combined impacts of wildlife harvesting by the community as a whole. Through the use of semi-structured interviews, we tested the hypothesis that larger communities with greater access to the economic market exert higher pressure on wildlife and target more commercially valuable species. These communities are expected to be affected by higher levels of wildlife depletion, with preferred species disappearing near villages, triggering shifts in harvested species spectra.

Study area

This study was carried out in the PSNR, which covers an area of 2,080,000 ha in the Department of Loreto, in the north-eastern Peruvian Amazon. It is bordered by two tributaries of the Amazon River, the Ucayali and Marañón rivers, and encompasses the two major drainage basins of the Pacaya and Samiria rivers. The reserve is characterised by massive hydrological fluctuations that occur between the high-water (October to May) and low-water (June to September) seasons (Kvist et al., 2001).

The majority of inhabitants are descendants of the Tupi-Guarani speaking Kukama-Kukamilla people and more recent immigrants of Caucasian and Indigenous origin (Gow, 2007). Their main livelihood activity is fishing, which is most productive during the low-water season, when fish become trapped in the shrinking water bodies. Nonetheless, migrations of fish feeding on fallen fruit in the *várzeas* (white-water flooded forests) make some fisheries productive during the high-water season (Kvist et al., 2001). The Kukama-Kukamilla also engage in opportunistic hunting, primarily during the high-water season, when the terrestrial fauna is concentrated on the non-inundated *restingas* (levees) (Bodmer et al., 1998).

Approximately 100,000 people live in over 200 communities along the boundary of the PSNR (INRENA, 2009). We selected five Kukama-Kukamilla villages located at the mouth of the Samiria River, which were divided into two distinct areas: a) San Martín de Tipishca, Nuevo Arica and Bolivar lie on the shores of the Tipishca Lake; and b) San José de Samiria and Leoncio Prado are located along the Marañón River (Fig. 1). These villages ranged from 40 to 120 households (Table 1), and differed in their exposure to the market economy. The communities of the Marañón River supply produce to the urban markets of Loreto by selling to freezer vessels or directly to market vendors.

Methods

Data collection

We conducted 122 semi-structured interviews, which accounted for 34.9% of

households within the study area, between June-August 2013 (Table 1). The use of semi-structured interviews was the preferred data collection method, as they allow emphasis on specific topics depending on the interviewees' knowledge and experience (Rubin & Rubin, 2005). Recall bias was expected to be minimal, as quantitative information asked was simple and activities are regular and highly seasonal (Golden et al., 2013). All households were found to be dependent on hunting and/or fishing, so we adopted a convenience sampling approach, selecting the most accessible households (Patton, 2002). We targeted male heads of households for interviews, but in some cases interviewed women instead, either because they too participated in hunting or fishing, or more often they had acquired detailed information about harvests through cooking. We obtained prior informed consent from participants before conducting interviews.

The social sensitivity of the topic being explored may have created some bias in the data resulting from the under-representation of harvests. Where possible, we used participant observation to verify interview responses. We informed interviewees that no information gathered would be used against them and that survey information would be anonymised.

Data analysis

We obtained household harvest rates of fish by asking fishermen to state the mean total biomass of fish caught per day, during high- and low-water seasons separately. This was extrapolated to annual harvest rates by multiplying each estimate of mean daily yield for each season by 182.5 (6 months). A limitation of using interviews to collect harvest data was that fishermen were unable to state the quantity of each species harvested, because

they measure the weight of the entire catch. We therefore recorded the percentage of households that harvest each species, using these data as proxies for relative harvest rates. We obtained annual household harvest rates of game species by asking hunters to state the mean number of wild animals hunted per year for each species, as hunting is less frequent than fishing. This was converted to biomass using body weight data reported by Peres and Dolman (2000), Ohl-Schacherer et al. (2007), Cardoso et al. (2012), and Mayor et al. (2015). We calculated per-capita harvest rates, assuming an average of six individuals per household. We determined total community-level harvest rates of fish by multiplying mean household harvest rates by the number of households in each community, and in the case of game species, by the percentage of households that engage in hunting.

We used household harvest rates to estimate catch-per-unit-effort (CPUE). The assumption behind CPUE as an indicator of sustainability is that hunters and fishermen must increase their efforts in areas with depleted populations to achieve the required meat and fish return rates. A difference in CPUE is assumed to reflect a difference in actual prey density or abundance (Rist et al., 2010). We calculated CPUE of fish as Y/H and CPUE of game species as B/D , where Y is the total daily yield of fish harvested; H is the number of hours a day fishermen leave their nets in the water (the most common method); B is the total biomass of games species hunted annually; and D is the number of days a year hunters are active. We averaged across households to obtain community-level CPUE estimates.

We calculated the distance travelled on hunting and fishing trips using reports of average time travelled. Based on information given by a local informant, we estimated

that 6 km were travelled in 1 hour in *peque peque* (motorized canoe) and 4 km on foot. Since hunters use watercourses to navigate to hunting sites and limit their activities to within 2 km into the forest from the river's edge, distance travelled was multiplied by four to obtain the size of the total catchment area (Begazo & Bodmer, 1998). The corresponding catchment area was drawn around the channels and lakes of the Samiria and Marañón rivers and divided into zones of low, medium, and heavy exploitation, using the maximum distances travelled by the top 25% and 50% percentiles as the thresholds (Fig. 2). Given our project's social science dimension and use of interviews, we determined that this measure of relative exploitation was appropriate (Brodizio & Chowdhury, 2010; Hawken & Munch, 2012). We used Welch's analysis of variance and the Kruskal-Wallis H test to compare distance travelled on hunting and fishing trips between communities. The Pearson's rank correlation coefficient allowed us to examine the relationship between CPUE and distance travelled as an indication of local resource depletion (Fa et al. 2006; Laurance et al. 2006).

We used multiple linear regressions to investigate the effects of socio-economic variables on CPUE and harvest rates. We included village size as a continuous variable and market exposure as a categorical variable in all models, using season as an additional categorical variable in the analyses of fishing data. The response variables were log-transformed to account for non-normal distributions. We estimated the significance of variables by dropping them from the full model and using likelihood ratio tests to compare nested models. We examined variations in the species compositions of harvests, termed the 'harvest profile', using Principal Components Analysis (PCA). Results were considered significant for $P < 0.05$. Statistical analyses were undertaken in R version 3.3.1 (R Core Team, 2016).

Results

All households in the study area fished daily throughout the year. In 57% of households, fishing was supplemented with hunting. 77% of hunters were active less than 10 days a year, and only one hunted as often as 18 days a year. The total biomass of wildlife harvested annually by the five communities was ~1,807 t (Table 1). The majority of fishermen (96%) reported travelling in *peque peque* for no more than 6 hours, whereas 39% of hunters undertook trips of several days, travelling over 6 hours to reach remote *restingas* inside the reserve. The mean distance travelled by fishermen and hunters was 11.2 (± 4.1) km and 44.0 (± 11.1) km, respectively. The distance travelled on hunting trips did not differ between communities ($H_{(4)} = 5.70$, $P = 0.22$), but fishermen from Nuevo Arica and San Martín de Tipishca travelled farther than fishermen from other villages (*Welch's* $F_{(4,29,67)} = 18.21$, $P < 0.001$). The combined hunting and fishing catchment area for all communities covered ~576 km² (Fig. 2). There was a positive correlation between distance travelled into the reserve and CPUE of fish during the low-water season (Pearson $r_{s(120)} = 0.22$, $P = 0.017$), but not the high-water season (Pearson $r_{s(120)} = 0.17$, $P = 0.07$). No significant correlation existed between distance travelled and CPUE of game species (Pearson $r_{s(69)} = 0.14$, $P = 0.24$).

The communities of the Samiria basin collectively harvested 1,740 t of fish annually (96.3% of biomass extracted), comprising 23 fish species (Table 2). The most widely caught species was *Prochilodus nigricans*, a species of both commercial and subsistence importance. There was substantial variation in harvest profiles between communities (Fig. 3). In San José de Samiria and Leoncio Prado, fishermen harvested a large

proportion of small, commercial species such as *Leporinus* spp., as well as larger species like *Hoplias malabaricus*. In San José de Samiria, smaller, less economically valuable species like *Oxydoras niger* and *Leiarius marmoratus* also made up a significant proportion of their catch. The communities of the Tipishca Lake depended on the most abundant species, including *Pterygoplichthys pardalis*, *Pygocentrus* spp. and *Serrasalmus* spp. We found evidence that the paiche, a species of conservation concern, was also caught.

The reported total annual harvest of game species in the study area was ~4,275 individuals, equating to ~67 t (3.7% of biomass extracted) and comprising 27 species (Table 3). Mammals were the most frequently extracted group, making up 74.0% of hunted biomass and 56.0% of all hunted individuals, followed by reptiles (23.1%; 19.1%) and birds (2.9%; 24.9%). The majority of biomass harvested came from large-bodied animals, mainly the white-lipped peccary (*Tayassu pecari*), lowland tapir, and black caiman *Melanosuchus niger*. The white-lipped peccary, paca *Cuniculus paca* and brown agouti *Dasyprocta variegata* were the most frequently hunted in terms of number of individuals. The Amazonian manatee *Trichechus inunguis*, which is strictly protected, was hunted occasionally. As with fish harvests, harvest profiles of game species varied substantially between communities (Fig. 4). In San José de Samiria and San Martín de Tipishca, hunters harvested a larger proportion of large-bodied species, such as the lowland tapir, the South American river turtle *Podocnemis expansa* and the white-lipped peccary, whereas the other communities harvested a larger proportion of small primates and wetland birds.

The multiple linear regressions revealed a significant positive relationship

between village size and annual community-level harvest rates of fish (Table 4, Fig. 5). However, village size had no effect on mean per-capita harvest rates ($F = 0.33$, $P = 0.59$) or CPUE ($F = 0.96$, $P = 0.37$) of fish. In contrast, there was no effect of market exposure on community-level harvest rates of fish ($F = 4.60$, $P = 0.08$), but commercial communities had significantly lower mean per-capita harvest rates and CPUE of fish (Table 4, Fig. 6-7). As expected, season had a significant effect on harvest rates and CPUE of fish, both of which were higher in the low-water season (Table 4). Neither market exposure nor village size had a significant effect on harvest rates or CPUE of game species (all $P > 0.31$).

Discussion

Our study adds to the growing body of research that suggests that socio-economic factors influence wildlife harvesting by Indigenous peoples (Smith & Wishnie, 2000; Lu, 2007; Godoy et al., 2010). Specifically, the patterns of hunting and fishing by the Kukama-Kukamilla people of the PSNR reveal the potential threat of increased market integration and a rising human population. The data presented in this study include a number of potential sources of variation that we did not control for, including environmental variables such as habitat quality, which may have limited the statistical power of the analyses. Furthermore, the small sample size of only five communities means caution must be taken when interpreting the results of the significance tests. However, because data points represent aggregates of household-level data, they reflect many more underlying observations, and we believe this allows us to make reliable inferences.

We discovered evidence that increased market exposure leads to resource depletion, reflected in a lower CPUE of fish in commercial communities. A reduction in fish populations as a result of over-fishing may have reduced the profitability of fishing and limited commercial fishing activity, which would explain why fishermen in commercial communities had lower mean per-capita harvest rates (Vasco & Sirén, 2016). Nevertheless, the net pressure that commercial fishing puts on depleted resources is likely greater than the pressure exerted by non-commercial communities on relatively un-depleted fish stocks. In San José de Samiria and Leoncio Prado, fishermen targeted small, economically valuable species, indicating possible over-exploitation of larger species. This trend is observed in the nearby markets of Iquitos, where the sale of cheaper, smaller and faster-growing species has risen since the 1980s and the sale of larger species has declined (Garcia et al., 2008; Atwood et al., 2015). The large proportion of less economically valuable species in harvests from San José de Samiria could reflect an increasing reliance on these species for subsistence.

As expected, larger communities exerted greater pressure on fish resources through increased harvest rates, because there is both more people to feed and a greater number of fishermen. We therefore expected to see similar signs of resource depletion in these communities. Nonetheless, community size had no significant effect on CPUE of fish. However, fishermen from San Martín de Tipishca, the largest village, together with those from Nuevo Arica, travelled farther on fishing trips than those from neighbouring communities, and during the low-water season CPUE was higher farther into the reserve. This is consistent with the paradigm that Neotropical people are central-place foragers, travelling greater distances in search of preferred prey species as wildlife populations become locally depleted (Levi et al., 2009; 2011). Thus, fishing in

previously un-exploited sites inside the PSNR could be masking resource depletion in the Tipishca Lake. Fishermen from San Martín de Tipishca also harvested small, abundant fish species, which may be able to sustain the larger human population.

We found no clear effect of village size or market exposure on harvest rates or CPUE of game species. This implies that people in larger, commercial villages have been able to shift to alternative sources of protein, such as fish or livestock, to meet subsistence and commercial needs. The strong presence of preferred species in harvest profiles suggests that wild meat harvests in the PSNR are currently supplied by a relatively un-depleted source. In San José de Samiria and San Martín de Tipishca, hunters harvested large-bodied prey species, including ungulates, large primates and reptiles. Encounter rates of these species in the forest are relatively low due to naturally low population densities, so hunters are likely targeting them for their greater meat harvests, as occurs in other Amazonian communities (Peres & Lake, 2003; Zapata-Ríos et al., 2009; Espinosa et al., 2014; Sirén & Wilkie, 2016). The current hunting patterns of the Kukama-Kukamilla people may be indicative of a source-sink dynamic, with immigration of game species from the un-hunted core zone of the reserve sustaining harvests in the catchment area (Navaro et al., 2000; Ohl-Schacherer et al., 2007).

Nevertheless, large-bodied game species are particularly vulnerable to over-exploitation due to slow reproductive rates (Mayor et al., 2017). The continued harvest of vulnerable species by larger, commercial communities will likely cause significant population declines in the PSNR and a shift in prey selection toward a broader range of smaller, less-preferred species, following the general trend observed throughout the Amazon (Naranjo & Bodmer, 2007; Peres & Palacios, 2007; Constantino, 2016). The

region has also been experiencing more extreme droughts and seasonal flooding in the last few decades, which could exacerbate the impacts of unsustainable wildlife extraction by limiting resources for wildlife and causing direct mortality of animals (Bodmer et al., 2017). The recent sharp decline in populations of the white-lipped peccary throughout its range, for which non-anthropogenic impacts are suspected, will put further pressure on alternative and more vulnerable prey species (Fragoso, 2004; Richard-Hansen et al., 2013; Mayor et al., 2015).

Overall, our results indicate that the forests of the PSNR are able to provide important food supplements for the Kukama-Kukamilla people. However, hunting and fishing in some villages appears to be approaching critical thresholds, threatening the natural capital of the reserve. Around the world, the combination of human population growth and increased market integration of Indigenous peoples is linked to a downward spiral of local species extinctions and a diminishing supply of crucial protein and income. In this context, the sustainable management of natural resources represents a crucial opportunity for biodiversity conservation where protected areas and Indigenous territories overlap (Zimmerman et al., 2001). Development professionals, protected area managers, and conservationists need to help maintain low hunting and fishing pressure by diversifying and enhancing existing livelihood strategies, thereby reducing poverty in rural communities and conserving vulnerable species (Bodmer & Lozano, 2001; Bassett, 2005; Gandiwa, 2011). Community-based management is needed to monitor the impacts of socio-economic and climatic change, and to ensure the long-term sustainable use of forest species, both inside and outside protected areas.

Author contributions

REB and MK designed the data collection methods and REB provided logistical support in the field. MK collected and analysed the data and wrote the first draft of the manuscript. JCA, MK, CE, REB and PM edited the manuscript to produce the final draft. AB produced the maps.

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Biographical sketches

Maire Kirkland conducts research into the sustainable use of natural resources. Cristina

576 Eisenberg works on food-web relationships, sustainable natural resources use, and
577 ecological restoration globally, with a focus on Indigenous communities. Andy Bicerra,
578 Richard E. Bodmer and Pedro Mayor are involved in wildlife research and biodiversity
579 conservation in the Neotropics. Jan C. Axmacher explores patterns of biodiversity in
580 China and the UK.

Table 1. Details of interviews and harvest rates in the five Kukama-Kukamilla communities located at the mouth of the Samiria River. The amount of meat available for consumption refers to the edible portion of fish and game meat, which was calculated as 70% of biomass extracted (Hill et al., 1984; Roos et al., 2007).

Community	San Martín de Tipishca	Nuevo Arica	Bolivar	Leoncio Prado	San José de Samiria
Number of families	120	50	40	90	50
Number interviewed (%)	29 (24.2%)	28 (56.0%)	9 (22.5%)	30 (33.3%)	26 (52.0%)
Total community-level harvest per year (t)					
Fish	679.64	222.26	359.32	327.95	151.26
Game	15.01	14.42	8.40	9.94	10.70
Total meat	694.65	236.68	367.72	337.89	161.96
Total harvest per-capita per year (t)					
Fish	0.94	0.74	1.50	0.61	0.51
Game	0.04	0.09	0.06	0.03	0.06
Total meat	0.98	0.83	1.56	0.64	0.57
Total meat available for consumption					
Per household per year (t)	4.11	3.47	6.55	2.68	2.39
Per-capita per year (t)	0.68	0.58	1.09	0.45	0.40
Per-capita per day (kg)	1.88	1.59	2.99	1.22	1.09

Table 2. Fish species harvested by the Kukama-Kukamilla people, showing the proportion of households harvesting each species during high- and low-water seasons.

Species			Percentage of households (%)	
Order	Scientific name	Local name	High	Low
Characiformes	<i>Prochilodus nigricans</i>	Boquichico	83.33	77.12
	<i>Hoplerethrinus unitaeniatus</i>	Shuyo	60.83	41.18
	<i>Mylossoma duriventre</i>	Palometa	26.67	45.00
	<i>Hoplias malabaricus</i>	Fasaco	26.67	28.57
	<i>Triportheus</i> spp.	Sardina	25.00	22.69
	<i>Leporinus</i> spp.	Lisa	15.83	22.69
	<i>Pygocentrus/Serrasalmus</i> spp.	Piraña	14.17	17.65
	<i>Potamorhina latior</i>	Yahuarachi	6.67	4.20
	<i>Brycon</i> spp.	Sabalo	5.83	5.74
	<i>Colossoma macropomum</i>	Gamitana	0.83	0.83
Perciformes	<i>Satanoperca jurupari</i>	Bujurqui vaso	15.00	23.33
	<i>Astronotus ocellatus</i>	Acarahuazú	9.17	26.27
	<i>Cichla monoculus</i>	Tucunaré	0.83	6.67
Siluriformes	<i>Pterygoplichthys pardalis</i>	Carachama	64.17	51.28
	<i>Pseudoplatystoma tigrinum</i>	Tigre zúngaro	4.17	5.83
	<i>Pimelodus blochii</i>	Bagre	3.33	4.17
	<i>Pseudoplatystoma fasciatum</i>	Doncella	3.33	2.50
	<i>Hoplosternum</i> spp.	Shirui	2.50	0.83
	<i>Hypophthalmus edentatus</i>	Maparate	0.83	1.67
	<i>Oxydoras niger</i>	Turushuqui	0.83	0.83
	<i>Leiarius marmoratus</i>	Achara	0.83	0.83
	<i>Sorubim lima</i>	Shiripira	0.83	0.00

Table 3. Annual per-capita harvest rates of game species by the Kukama-Kukamilla people, showing the biomass and number of individuals harvested per person per year.

Species			Annual per-capita harvest	
Order	Scientific name	Local name	Biomass (kg)	Number of individuals
MAMMALIA				
Artiodactyla	<i>Tayassu pecari</i>	Huangana	17.33	0.50
	<i>Pecari tajacu</i>	Sajino	2.60	0.10
	<i>Mazama americana</i>	Venado colorado	1.02	0.05
Perissodactyla	<i>Tapirus terrestris</i>	Sachavaca	13.19	0.10
Rodentia	<i>Cuniculus paca</i>	Majáz	3.38	0.42
	<i>Dasyprocta variegata</i>	Añuje	1.63	0.33
Sirenia	<i>Trichechus inunguis</i>	Vaca marina	2.90	0.01
Cingulata	<i>Dasyus novemcinctus</i>	Carachupa	1.61	0.27
Primates	<i>Alouatta seniculus</i>	Coto	1.19	0.18
	<i>Sapajus apella</i>	Mono negro	0.25	0.09
	<i>Cebus albifrons</i>	Mono blanco	0.19	0.07
	<i>Saimiri boliviensis</i>	Maquisapa	0.18	0.02
	<i>Lagothrix</i> spp.	Choro	0.04	0.01
	<i>Ateles</i> spp.	Fraile	0.01	0.02
Carnivora	<i>Nasua nasua</i>	Achuni	0.17	0.03
REPTILIA				
Crocodilia	<i>Melanosuchus niger</i>	Lagarto negro	7.00	0.15
	<i>Caiman crocodilus</i>	Lagarto blanco	1.60	0.05
Testudinae	<i>Podocnemis unifilis</i>	Taricaya	2.78	0.35
	<i>Chelonoidis denticulata</i>	Motelo	1.02	0.13
	<i>Podocnemis expansa</i>	Charapa	1.88	0.07
AVES				
Anseriformes	<i>Cairina moschata</i>	Sachapato	0.58	0.19
Galliformes	<i>Mitu tuberosum</i>	Paujil	0.29	0.10
	<i>Pipile cumanensis</i>	Pava	0.22	0.16
	<i>Penelope jacquacu</i>	Pucacunga	0.15	0.12
Pelecaniformes	<i>Phalacrocorax brasilianus</i>	Cushuri	0.25	0.17
	<i>Ardea</i> spp.	Garza	0.24	0.20
Tinamiformes	<i>Crypturellus undulatus</i>	Panguana	0.04	0.04

Table 4. Results of the multiple linear regression analyses showing the effect of village size, market exposure and season on log-transformed harvest rates and CPUE. Non-significant variables were excluded from each model.

	Estimate \pm SE	<i>t</i> -value	<i>P</i> -value
Community-level harvest rates			
(Intercept)	3.41 \pm 0.35	9.74	<0.000
Village size	0.01 \pm 0.00	2.66	0.032
Low-water season	1.25 \pm 0.26	4.80	0.002
Per-capita harvest rates			
(Intercept)	-1.52 \pm 0.18	-8.61	<0.000
Commercial	-0.54 \pm 0.22	-2.44	0.045
Low-water season	1.25 \pm 0.22	5.76	<0.001
CPUE			
(Intercept)	-0.80 \pm 0.14	5.73	<0.001
Commercial	-1.10 \pm 0.17	-6.27	<0.001
Low-water season	1.61 \pm 0.17	6.78	<0.001